

Doppler optical coherence tomography for interventional cardiovascular guidance: *in vivo* feasibility and forward-viewing probe flow phantom demonstration

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Abstract. We demonstrate the potential of a forward-looking Doppler optical coherence tomography (OCT) probe for color flow imaging in several commonly seen narrowed artery morphologies. As a proof of concept, we present imaging results of a surgically exposed thrombotic occlusion model that was imaged superficially to demonstrate that Doppler OCT can identify flow within the recanalization channels of a blocked artery. We present Doppler OCT images in which the flow is nearly antiparallel to the imaging direction. These images are acquired using a flexible 2.2-mm-diam catheter that used electrostatic actuation to scan up to 30 deg ahead of the distal end. Doppler OCT images of physiologically relevant flow phantoms consisting of small channels and tapered entrance geometries are demonstrated. © 2010 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.3292007]

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1 Introduction

Chronic total occlusions (CTOs) are defined as arterial blockages that appear occluded under x-ray angiography and are older than 12 weeks.¹ Patients that have a CTO are typically sent for coronary artery bypass (open heart) surgery instead of minimally invasive intravascular therapy such as angioplasty. The principal challenge of performing angioplasty for a CTO is the safe navigation of a guidewire through the dense collagen blockage without perforating the arterial wall.^{2,3} While some experimental imaging guidance technologies have been suggested to differentiate between the blockage and the arterial wall,⁴ and hence guide this procedure, these methods are not generally available to interventional cardiologists.

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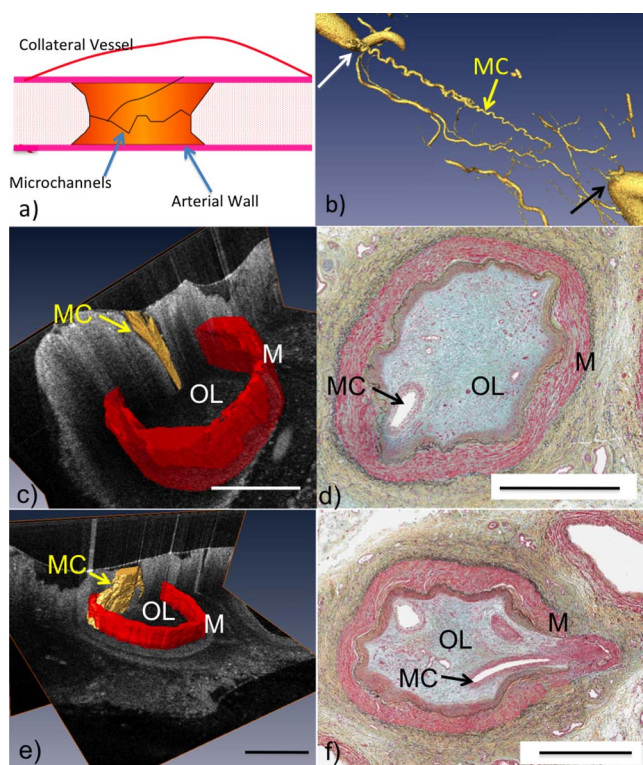


Fig. 1 Microchannels in CTOs. (a) Drawing representing a CTO illustrates both a microchannel that traverses the lesion and one that exits through the arterial wall. (b) Parallel studies with x-ray microcomputed tomography of rabbit arterial occlusion illustrate that these occlusions do possess long channels (labeled as MC) through the lesion. The white arrow in (b) identifies the proximal entrance, while the black arrow shows the location of the distal end. (c) A 3-D OCT volumetric image of a segment of an arterial occlusion, where the arterial medial wall (M) was manually segmented as red and a traversing microchannel (MC) as yellow and (d) corresponding Movat-stained histology. (e) A Volumetric OCT image of an arterial occlusion in which a microchannel appears to exit through the arterial wall and (f) the corresponding Movat histology. (Color online only.)

Despite their name, however, more than half of CTOs are actually less than 95% occluded under histological observation.⁵ Small channels are routinely observed in these lesions, and these channels may either remain within the lumen or exit the lesion, as shown schematically in Fig. 1(a). Parallel studies with x-ray micro computed tomography have shown that these channels traverse a significant distance through the lesion, as shown in Fig. 1(b). We previously postulated that these channels may serve as useful conduits for a guidewire to follow through the occluded lumen⁶ as they may represent the path of least resistance through the occlusion. However, it has also been observed in histology that these channels occasionally exit from the lumen,⁵ potentially representing dangerous points where it would be easier to rupture the arterial wall. These channels are typically too small⁵ (50 to 200 μm) to be visualized using x-ray angiography and therefore present an attractive imaging target for a high-resolution intravascular imaging technique.

Intravascular imaging using optical coherence tomography (OCT) has attracted a great deal of attention for diagnosing arterial plaques that are prone to rupture.⁷⁻⁹ In previous work,

we demonstrated that a time-domain OCT system can identify different pathologies *ex vivo* that are observed in CTOs in the peripheral arteries.¹⁰ We also were able to expand on this work to include 3-D volumes of human *ex vivo* occluded artery segments acquired from peripheral limb amputations using a swept source OCT system. In Figs. 1(c) and 1(e) we show two examples of 3-D OCT images in which the arterial wall (tunica media) and microchannels were manually segmented based on their optical backscattering properties. In the first example [Fig. 1(c)], we were able to identify a channel that appears to pass directly through the occlusion, while in the second, [Fig. 1(e)], we observed a channel that appears to exit out of the media wall. The corresponding histology is shown in Figs. 1(d) and 1(f).

While these *ex vivo* imaging results are encouraging, *in vivo* imaging identification of these microchannels based solely on differences in reflectivity of the occlusive collagen and the blood-filled microchannel may prove difficult as both blood and collagen have high backscattering properties. One additional contrast mechanism for identifying channels is the use of flow information. Flow may also better indicate continuous channels. Previous high-frequency ultrasound imaging work by Thind et al. was able to identify flow in the microchannels of a pig femoral artery CTO model.¹¹

We therefore sought to demonstrate the potential of Doppler OCT as a means of providing contrast in a forward-viewing imaging geometry. The Doppler signal is defined by

$$f_d = \langle v \rangle 2n_t (\cos \theta) \lambda_0^{-1},$$

where $\langle v \rangle$ is the mean velocity of the pixel being imaged, n_t is the index of refraction of the object being imaged, θ is the Doppler angle (the angle between the imaging beam and the direction of the moving object), λ_0 is the center wavelength of the imaging light source, and f_d is the resulting Doppler shift. Therefore, the imaging geometry in which the flow direction is either parallel or antiparallel ($\theta=0, \pi$) to the imaging direction represents an optimal sensitivity to the Doppler frequency shift.

As a first step to evaluate the feasibility of Doppler OCT to detect flow in nearly occluded arteries, a surgically exposed occluded femoral artery was imaged from the surface with bulk optics in a 2-week-old rabbit model.

We then present work using a previously developed forward-viewing OCT catheter to image flow in phantoms that mimic nearly occluded lesions.

2 Methods and Materials

2.1 Occluded Artery Model

To test Doppler OCT's ability to detect *in vivo* flow within microchannels, we used a thrombotic occlusion model in the rabbit femoral artery that was previously described.^{12,13} Briefly, this model involves surgically exposing and isolating a rabbit femoral artery, ligating the artery both distally and proximally, injecting approximately 0.1 ml thrombin solution (200 international units/ml) at the proximal end of the artery and then removing the proximal ligature to allow blood to flow into the lesion and create a thrombus [as shown in Fig. 2(a)]. After approximately 1 h, the wound was closed and left for 2 weeks.

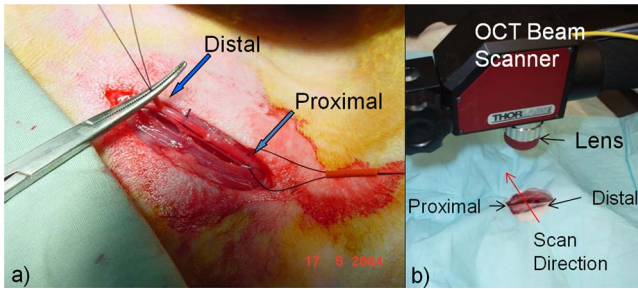


Fig. 2 (a) Creation of a rabbit thrombus occlusion, showing an isolated rabbit femoral artery with a closed ligature at the distal end and (b) image illustrating the scanning setup used for *in vivo* imaging of the exposed artery.

2.2 Intravital OCT Imaging

After 2 weeks, the wound was reopened and the artery was isolated. It was imaged using a Thorlabs OCT system (SL1325-P16, New Jersey, USA, center wavelength $\lambda_0 = 1325$ nm, bandwidth $\Delta\lambda = 110$ nm, and axial scan speed of 8 kHz) equipped with a handheld scanner attached to a surgical arm. The scanner consisted of a pair of galvomirrors and a lens, which scanned and focused the sample arm beam of the OCT system, respectively. This scanner was positioned above the artery, as shown in Fig. 2(b), such that it imaged across the artery creating a cross-sectional image of the occluded vessel.

2.3 Forward-Viewing Optical Probe

For flow phantom measurements (and eventual *in vivo* use), a forward-viewing scanning probe was constructed similar to that described previously.¹⁴ Briefly, the probe used a high-voltage, low-current electrode and a cantilever coupled to a ground electrode through a dissipative polymer to scan an optical fiber across a graded-index (GRIN) lens. For the experiments presented here, the probe was driven at 2200 V with oil immersion, which enabled scanning at 15 Hz (or 30 frames/s when both slopes of the triangular waveform are used for imaging). To effectively visualize the entire cross section of the occlusion a 20-mm working distance GRIN lens was used. This probe was packaged in a 2.2-mm-diam Teflon tube. A schematic of the probe layout is shown in Fig. 3(a) and photographs of the probe when stationary and in motion are shown in Figs. 3(b) and 3(c), respectively.

2.4 Fourier-Domain Mode-Locked OCT System

The forward-looking probe was connected to the sample arm of the OCT system, consisting of a custom-made swept-source OCT system based on a tunable laser with a center wavelength of 1320 nm and a repetition rate of 43 kHz. The laser employed¹⁵ the Fourier domain mode-locking technique to achieve a ranging depth of 6 mm. For *in vivo* use, the longer ranging depth may provide a larger field of view, given the relatively small sector scan angles provided by the forward-viewing probe.

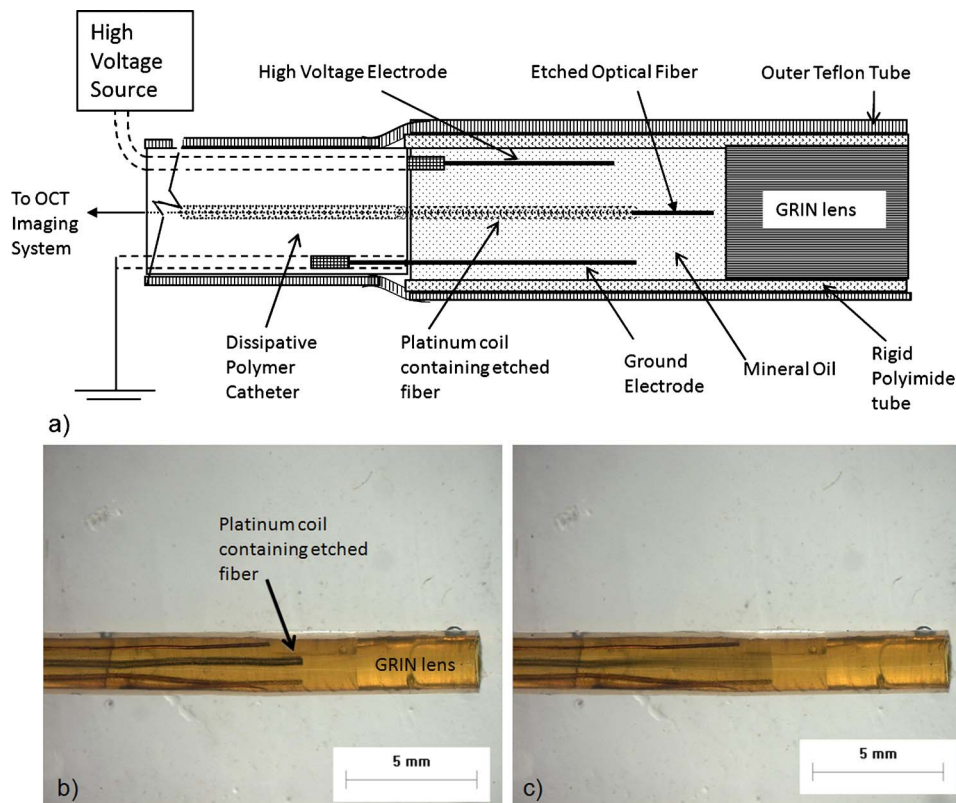


Fig. 3 Forward-viewing OCT scanning catheter: (a) a schematic of the probe showing the components used, (b) a photograph of the probe without the application of high voltage, and (c) the probe in oscillation, being driven by 2200 V.

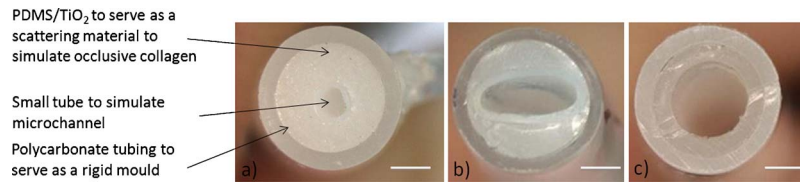


Fig. 4 Variety of phantoms that were created for imaging flow in the forward direction: (a) a cleaved end face of 600- μm -ID, 900- μm -OD tube embedded in titanium dioxide/PDMS to mimic the appearance of a central channel within an arterial occlusion (a polycarbonate tube is used as a rigid mould), (b) an oval-shaped channel that is used to mimic the appearance of a narrowed lesion, and (c) a pipette tip placed inside the polycarbonate tube to mimic a lesion that gradually tapers to a small channel. Scale bars represent 1 mm.

For OCT structural imaging, this system was able to generate 350 lines/mm when the scanning probe was operated at 15 frames/s. For Doppler signal processing, an ensemble length of 32 lines was used to generate the forward-looking Doppler images.

2.5 Flow-Phantom Construction

To create phantoms mimicking blunt occlusions with microchannels, a 600- μm -inner-diam (ID), 900- μm -outer-diam (OD) Teflon tube was placed inside a larger 2.8-mm-ID, 3.8-mm-OD polycarbonate tube such that the smaller tube extended significantly out from both ends of the larger tube. The 3.8-mm tube was then filled with a mixture of TiO_2 powder and polydimethylsiloxane (PDMS) at a ratio of 0.2 mg/ml and allowed to set. After the PDMS had cured, one end of the tube assembly was cut with a scalpel to create a clean face, as shown in the Fig. 4(a). This assembly was then inserted into a stretchable Tygon tube such that the Tygon tube fit snugly over the tube assembly to create a layered structure that allows for fluid to be injected at one end. Other

types of structures that were also created, including an oval-shaped channel [Fig. 4(b)] and a tapered entrance created using a pipette tip [Fig. 4(c)].

As a first step, the entire flow phantom and the distal tube were filled with saline to create an optically clear field of view. A 1% mixture of Intralipid and saline was then added to a 10-ml syringe, and this syringe was attached to the distal end of the small Teflon tube. The syringe was placed in an infusion pump to simulate flow coming through the microchannel. The probe was then placed inside the phantom and the high-voltage power supply that was connected to the probe was activated so that the probe began to oscillate. The distance from the probe to the cleaved face of the flow phantom was adjusted such that the entire proximal end of the “occlusion” could be visualized. The setup for this work is detailed schematically in Fig. 5.

Once a clear view of the proximal end of the flow phantom was obtained, the infusion pump was activated at rate of 1 ml/min. The data acquisition was initiated once the Intralipid was seen in the microchannel tube, several centimeters away from the imaging area. The purpose of this timing

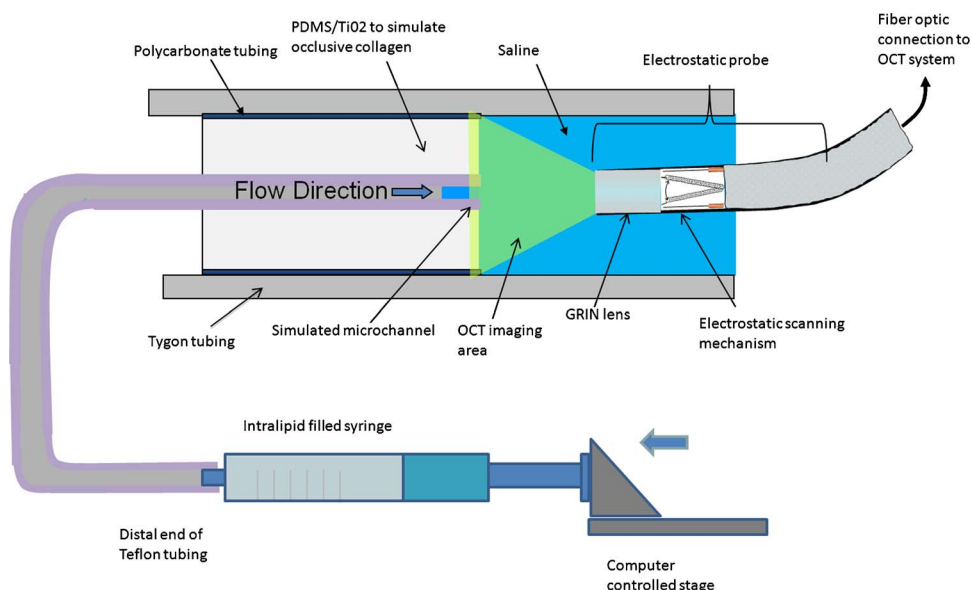


Fig. 5 Setup used for forward-viewing Doppler OCT in which the electrostatic probe images flow directed toward it through a small-diameter Teflon tube.

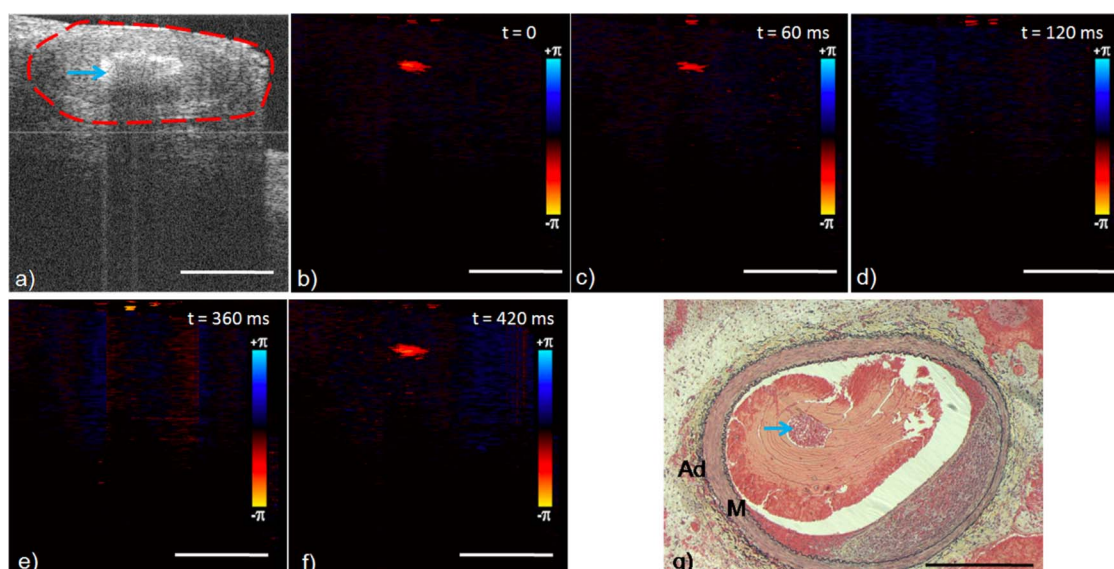


Fig. 6 Surgically exposed 2-week-old occlusion with both (a) structural and (b) to (f) a time sequence of Doppler images separated by 60 ms. The Doppler image identifies a central microchannel. A red dotted curve is used to identify the outer border of the artery on the structural image in (a). A representative Movat-stained histology slide is shown in (g). The adventitia is labeled Ad and the media is labeled M. Blue arrows identify the central microchannel in both the structural OCT (a) and histology images (g). Scale bars represent 0.5 mm. (Color online only.)

sequence was to capture the arriving bolus of Intralipid.

3 Results

3.1 *In Vivo DOCT Imaging*

Two-week-old arterial occlusions in a rabbit were exposed surgically and imaged. The structural OCT image in Fig. 6(a) shows the artery as outlined by the red dotted line. Within the artery is what appears to be a microchannel, as identified by the arrow. The Doppler OCT image sequence [Figs. 6(b)–6(f)] demonstrates that the central microchannel does possess pulsatile flow. The Movat-stained histology slide shown in Fig. 6(g) also illustrates the central microchannel within the occluded lumen. Using an estimated Doppler angle of 80 deg

and the known properties of the swept-source laser, we estimated the peak velocity seen in this channel as approximately 8 mm/s.

3.2 *Forward-Viewing Catheter Doppler OCT Imaging*

A time sequence of structural OCT images displaying the arrival of a bolus of Intralipid in the phantom is shown in Figs. 7(a)–7(c), with the corresponding forward-looking Doppler OCT (DOTC) images shown in Figs. 7(d)–7(f). The flow profile, as seen in the DOCT images [Figs. 7(d)–7(f)] appears parabolic through the channel. The Doppler images displayed aliased flow, as seen by the transition from yellow to blue (representing a transition in phase shift from negative to posi-

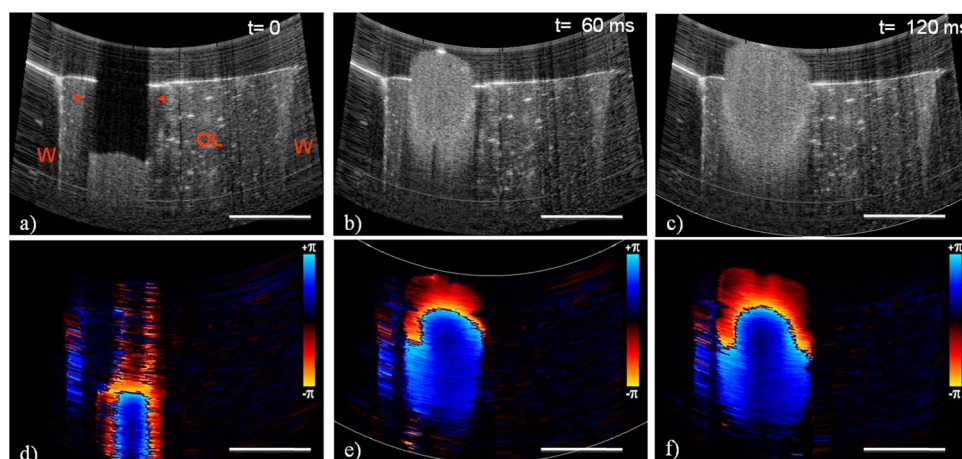


Fig. 7 Sequence of images taken every 60 ms showing both the structural (a) to (c) and (d) to (f) the corresponding Doppler progression of a bolus of Intralipid through a microchannel in an occlusion phantom. The occluded lumen composed of PDMS and TiO₂ is identified in (a) as OL. The Tygon™ tubing walls are seen as dark borders on either side of each image and are labeled with a W. Red arrowheads identify the outer walls of the embedded tube used to form the microchannel in (a). Scale bars represent 1 mm. (Color online only.)

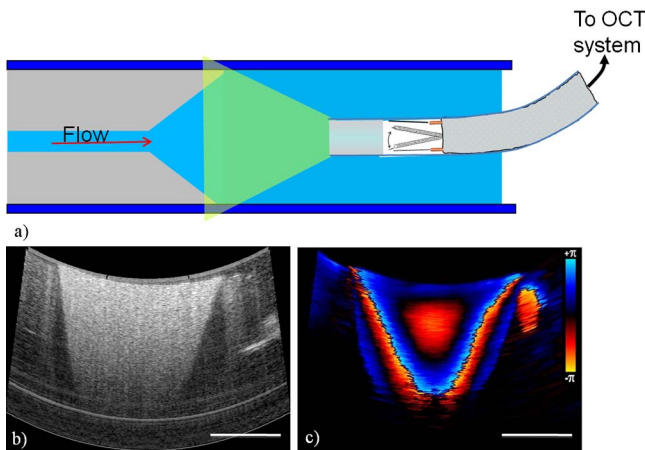


Fig. 8 (a) Schematic drawing of flow from a tapered exit occlusion directed toward the forward-viewing electrostatic probe (the red arrow indicates the direction of flow), and (b) structural and (c) Doppler images of Intralipid flow arriving in a tapered phantom [as shown in Fig. 4(c)] are displayed. Scale bars in (b) and (c) represent 1 mm. (Color online only.)

tive π). In this situation, the phase difference between the adjacent lines is greater than 2π . However, the measurement range for phase is limited from $-\pi$ to π , corresponding to ~ 11 mm/s. Thus, phase differences greater than this range will jump between $-\pi$ and π .

We were also able to examine several other geometries of the CTO phantoms, including imaging flow through a tapered entrance, as shown in Fig. 8, and the oval-shaped microchannel, as shown in Fig. 9. In both cases, the Doppler images serve to provide additional contrast over the structural images. This contrast is important in situations where the surrounding occlusion and the microchannel fluid have similar scattering properties, such as with collagen and blood.

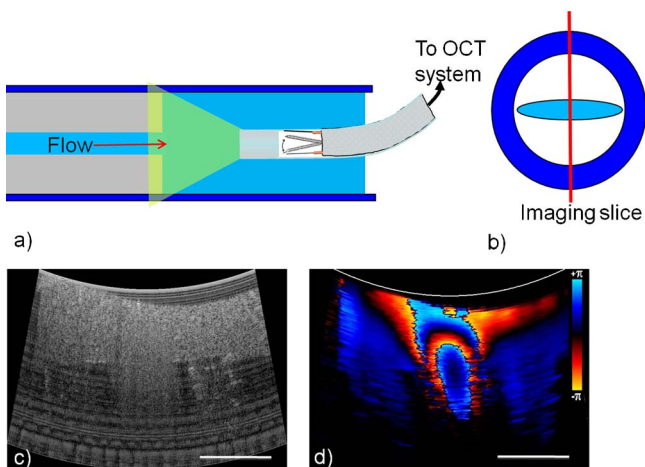


Fig. 9 Imaging flow through an oval channel using a forward-viewing Doppler OCT probe: (a) the orientation of the probe with respect to the flow phantom [as shown in Fig. 4(b)], (b) the face-on schematic of the flow phantom and the orientation of the imaging slice through it, (c) the structural image of Intralipid arriving through the channel, and (d) the corresponding Doppler image. Scale bars in (c) and (d) represent 1 mm.

4 Discussion

Advanced real-time imaging techniques in interventional cardiology capable of guiding procedures in arteries that appear occluded under x-ray angiography remain a highly desired but unrealized goal. Success rates for opening such arteries, in both the peripheral and coronary circulation, are much lower than for stenotic but not fully occluded lesions,¹⁶ in large part due to inadequate visualization of the occluded segment.

This preliminary study demonstrates that Doppler OCT can image flow in the small channels that recanalize an arterial occlusion. The calculated flow velocity in the channel detected in a 2-week-old occlusion is similar in value to the Doppler ultrasound results in microchannels of a pig femoral arterial occlusion.¹¹ This study also demonstrates Doppler imaging in flow phantoms with a forward-viewing catheter in which the imaging direction and the flow direction were nearly antiparallel. The Doppler images provide an added contrast mechanism that may help to identify small channels and the geometry of the lumen. This information in turn may help to define the overall morphology of the occlusion as well as identify points where the channels exit the occlusion.

This study was limited in its intravascular application to the imaging of an arriving bolus of Intralipid in flow phantoms. This limitation was a result of the small angular field of view of the probe and the requirement of imaging an entire cross section of the flow phantom. Reconciling these two points required the use of a longer working distance lens in the probe. However, because of the limited imaging penetration of OCT through Intralipid, this long working distance made imaging cases of constant Intralipid flow severely restricted. Thus, we were only able to study cases in which flow was directed toward the imaging catheter. In an *in vivo* implementation, flow would likely be directed away from the catheter either by pulsatile blood flow or through the injection of a blood and saline mixture to obtain an optimal balance of backscattered signal and imaging depth. In such an imaging geometry, it is likely that the probe itself will alter the flow pattern. Investigation of these effects remains a subject of ongoing endeavors.

Practical *in vivo* implementation requires several important improvements to the current probe design. Increasing the angular field of view is necessary to enable the probe to be placed closer to the proximal end of the occlusion while maintaining a full visualization of both the blockage and the two sides of the arterial wall. Furthermore, to generate 3-D forward-looking images analogous to those shown in Figs. 1(c) and 1(e) *in vivo*, image acquisition time would need to be improved by a factor of 20. Possible methods to implement such an increase include moving toward a fiber bundle/array design¹⁷ or a fast-scanning piezoelectric oscillator design.¹⁸ The probe's dimensions also require considerable miniaturization to be used effectively *in vivo*. The probe used in this work had a diameter of 2.2 mm with a rigid length of 15 mm. For use in a clinically relevant imaging scenario in interventional cardiology, a diameter of 1.0 mm with a rigid length of less than 4 mm would be required. To address this miniaturization issue, we are currently investigating methods of incorporating planar electrodes directly into the catheter's outer sheath as well as investigating fiber array designs, which may be more amenable to miniaturization. Development of an effective

means to clear the blood field to enable for imaging of the proximal end of the occlusion will also require careful thought. Possible methods include the incorporation of a balloon onto the imaging catheter or the use of saline flushing ports directly incorporated onto the outer sheath of the catheter. Theoretically the relatively small blood flow volume through the microchannel(s) in CTO lesions may reduce the difficulty of saline flush compared to conventional coronary OCT imaging requirements. Integrating therapeutic means with a forward-looking OCT catheter will also be necessary for this technology to be adopted in a clinical environment. Incorporating laser ablation techniques via optical fibers embedded within the outer sheath of the imaging catheter is a particularly attractive solution as it may enable simultaneously imaging and ablating the blockage. The use of ultrafast laser ablation may also be advantageous, as this method would enable the removal of calcified tissue.¹⁹

Despite these challenges, the ability to image the proximal end of the occlusion and to identify the geometry of severely narrowed arterial lesions using Doppler OCT would provide a valuable tool for interventional cardiology. The potential application using Doppler OCT data to guide wires through the lesion and to identify points where channels exit the vessel could make interventions on occluded arteries safer, more successful, and less invasive.

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